APPLICATION FOR LETTERS PATENT

Title: PEPTIDE LIGANDS THAT BIND TO SURFACES OF BACTERIAL SPORES

This invention was made with government support under the Defense Advance Research Project Agency project number MDA972-96-0003. The government of the United States has certain rights in the invention.

This application takes priority from Provisional Patent Application 60/071,411 filed January 14, 1998.

Background of the Invention:

The capture and identification of bacterial spores is useful for detecting pathogenic or otherwise harmful bacteria. Often the presence of spores can indicate to the researcher or epidemiologist the presence of virulent organisms. It is also important to determine the presence of spores of pathogenic organisms in the environment in order to more effectively control spread of infections. The ability to produce a monitorable tag or ligand that will bind specifically to the bacterial spore would provide a valuable tool for identifying pathogenic organisms in the infected patient and in the environment.

The use of phage-displayed peptide libraries to identify peptide sequences that will bind to particular receptors has been used to evaluate the structure of proteins. (See D'Mello, et al, <u>Virology</u>, **237**(2): 319-26 (1997) and Salonen, et al, Journal of General Virology 79 (pt4): 659-65 (1998) regarding mapping of antibodies and Marzari, et al, FEBS Lett. 411(1) 27-31 (1997) regarding phage display of B. thuringiensis insecti-A synopsis regarding the use of phage display cidal toxin.) has been reviewed by Smith and Petrenko in Chemical Reviews, 97(2) 391-410 (1997). In that review, the authors discuss the general usefulness of this procedure for evaluation of proteins, including antibodies. However, none of the above references suggest using phage display peptide libraries for identifying peptide sequences which bind to whole cells.

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Summary of the Invention:

This invention relates to the capture and/or identification of microorganisms and their spores by preparing peptides which bind to the spores. The peptides are included in phage display peptide libraries that are commercially available, and the peptides that bind to spores are identified using biopanning. While peptide sequences which bind to proteins, especially antibodies, have been studied, the method has not been used to identify peptides that bind to whole microorganisms.

The peptides of the invention and the methods by which the useful peptides are identified are disclosed herein. It is possible, using the peptides which bind to the surface of a cell which are generated by methods described herein, to identify the presence of spores of organisms in the environment and in the clinical setting. Using means of the invention, it is also possible to provide means for protecting potential hosts from exposure to disease-causing spores by administration of peptides which bind to the spores.

The peptides of the invention which bind to <u>Bacillus</u> <u>subtilis</u> contain the amino acid sequence Asn-His-Phe-Leu (NHFL) (Seq. ID No. 1). Additional amino acids containing proline, to provide the sequence NHFLP (Seq. ID No. 39) are particularly preferred sequences.

Peptides of the invention which bind to <u>Bacillus anthracis</u> have the sequence Thr-Ser-Gln-Asn-Val-Arg-Thr (TSQNVRT) (Seq. ID No. 40) or of the general formula Thr-Tyr-Pro-X-Pro-X-Arg (TYPXPXR) wherein X is a hydrophobic residue. Preferred residues are of the sequence Thr-Tyr-Pro-Ile-Pro-Ile-Arg (TYPIPIR) (Seq. ID No. 41), Thr-Tyr-Pro-Ile-Pro-Phe-Arg (TYPIPFR) (Seq. ID No. 42), and Thr-Tyr-Pro-Val-Pro-His-Arg (TYPVPHR) (Seq. ID No. 43).

Peptides which bind <u>Bacillus cereus</u> having sequences Val-Thr-Ser-Arg-Gly-Asn-Val (VTSRGNV) (Seq. ID No. 100) and consensus peptides of the formula Ser-Pro-Leu- X_1 - X_2 -His wherein X_1 is His or Arg and X_2 is Arg or Lys (SPLX₁X₂H) were also identified.

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The DNA sequence coding for the exemplified peptides is shown. It is clear that other codons that code for the same amino acid may be substituted using codon tables provided in molecular biology treatises.

The inventive method requires mixing phage from the phage display library with spores, incubating the mixture at about room temperature and separating the phage-spore complexes by centrifugation. The phage-spore complexes are washed several The phage was then eluted from the phagetimes in buffer. spore complexes with cold buffer at low pH, then quickly neutralized to prevent phage killing. The phage can then be amplified by infecting an appropriate organism. (E. coli was used in the instant case.) The cell lysate obtained from the culture may then be subjected to previous steps repeatedly. After about four rounds, individual clones are purified from Phage plaques (about 30 were used) were the eluted phage. amplified, the genomic DNA extracted and the DNA sequence of the 7-mer and 12- mer peptides encoding region determined. This DNA sequence indicates the sequence of the tight-binding The indicated protein sequences are tagged and exposed to known spores to determine binding properties.

Detailed Description of the Invention:

It is the purpose of this invention to identify short peptide ligands that bind specifically to the spores microorganisms, particularly those of Bacillus species. peptide ligands will bind tightly and in a species-specific manner to a physiological or fortuitous receptor on the surface of the spore. This peptide ligand can be used to capture the cognate spore in filters or as part of a detection device (e.g., a capture devise that concentrates the spores for identification by mass spectroscopy, DNA/RNA sequence evaluation, etc.) The peptide ligands can also be used directly in detection/identification devices and procedures. The peptides can be coupled to detectable (e.g., fluorescent, phorescent, radioactive, etc.) tags and the peptide-tag conjugates mixed with a sample which can contain cognate If spores are present, they will be bound by the spores.

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Spores of primary interest in the studies disclosed herein were spores of <u>Bacillus subtilis</u>, <u>Bacillus antracis</u>, <u>Bacillus</u> cereus, and Bacillus thuringiensis. However, methods of the invention may also be used to identify and capture other pathogens such as Clostridia species. B subtilis is a target primarily because it is used as a simulant in the development of detection devices for pathogenic B. anthracis. B. antracis is a key target because of its potential as an agent for use in biological warfare and terrorism. B. cereus and B. thuringiensis are targets because they closely resemble B. anthracis and because they are widely distributed in the environment. Thus, they can, potentially, produce false positive readings in detection devices and systems used to identify B. anthracis spores.

Phage Display ligand screening was employed using a commercially available (New England BioLabs) combinatorial library of 2 X 109 random peptide sequences (7-mer and 12-mer peptides were studied) were individually displayed on the surface of the filamentous coliphage M13. The random peptides were fused to the amino terminus of the minor coat protein The library is made by inserting a random nucleotide sequence at the beginning of the pIII gene of many copies of the M13 genome. These recombinant genomes are used to produce M13 phage. Each recombinant pIII gene produced a random peptide-pIII fusion protein and five copies of this fusion protein are displayed at one end of the mature phage particle. Thus, the random peptide sequence is displayed at the amino terminus of each pIII copy for a given phage. Furthermore, the random peptide sequence displayed by a particular phage clone can be readily determined by sequencing the peptide-encoding region of the phage genome.

Variations in the process would be known to one of skill in the art. Some of the modifications which enhance productiv-

ity are provided herein.

While the methods of the invention were first practiced targeting <u>B. subtilis</u>, then targeting <u>B. anthracis</u>, <u>Bacillus thuringiensis</u> and <u>B. cereus</u>, the methods disclosed herein, particularly the biopanning methods, may be used to identify useful sequences for binding to surfaces of other microorganisms.

The peptides of the invention may be prepared by means known in the art. These peptides can be synthesized, for example, using solid-phase synthesis and standard F-MOC chemistry. Then one would screen the compound using the methods described herein and comparable methods known in the art. Materials and Methods:

Peptides of interest were identified using a phage display ligand screening system. A phage display peptide library kit (New England BioLabs) was used according to instructions of the manufacturer in the identification process. The phage display library contains random 7-mer peptides (2 x 10 sequences) fused to the minor coat protein (pIII) of the filamentous coliphage M13. The phage containing the peptide ligands of interest were isolated from the phage library by several cycles of biopanning. The ligands of interest were then identified by sequencing the appropriate genomic region of the isolated phage.

In the biopanning procedure 10¹¹ phage were mixed with 10⁹ spores, incubated for 10 minutes at room temperature, and phage-spore complexes were separated by centrifugation at 4°C. The complexes were washed ten times with ice-cold TBST [50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 0.5% Tween 20] and phage were eluted from the complex with ice-cold elution buffer (0.2 m glycine-HCl, pH 2.2). The phage-containing eluate is immediately adjusted to a pH of about 7. This phage population is amplified by infecting E. coli, and the resulting lysate is used for a second round of biopanning. After about four rounds of biopanning, the eluted phage were plated to obtain single plaques, which were used to prepare large amounts of the particular phage. DNA was extracted from each phage prepara-

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tion, and the genomic sequence encoding the 7-mer peptide was determined. The DNA of over thirty independent phage isolates of <u>B. subtilis</u> were sequenced and thirteen unique sequences were identified. (See table 1.) All encoded peptides contained amino terminal sequence Asn-His-Phe-Leu. Although the sequences at positions five through seven are not identical, there is a clear preference for certain amino acids.

TABLE 1. Nucleotide and Amino Acid Sequences from <u>B. sub-</u>
<u>tilis</u> Spore-Binding Phage

| | | <u> </u> | <u>.s</u> spc | Te-D1 | rnarně | Phas | je | | |
|------------------|---------|----------|---------------|----------------------------------|--------|------|-----|----------------|------------------|
| 10 | Isolate | | | | | | | | |
| | 1 | AAT | CAT | TTT | TTG | ATT | AAG | CCG | (Seq. ID No. 2) |
| | 2 | AAT | CAT | TTT | TTG | AGG | TCT | CCG | (Seq. ID No. 3) |
| # | 3 | AAT | CAT | $\mathbf{T}\mathbf{T}\mathbf{T}$ | CTG | CCT | CGT | TGG | (Seq. ID No. 4) |
| | 4 | AAT | CAT | TTT | CTT | CCT | AAG | GTG | (Seq. ID No. 5) |
| N 15 | 5 | AAT | CAT | TTT | CTG | TTG | CCG | CCG | (Seq. ID No. 6) |
| 15 | 6 | AAT | CAT | TTT | TTG | CCT | CCT | CGG | (Seq. ID No. 7) |
| ≒ | 7 | AAT | CAT | TTT | CTG | CCT | ACT | GGG | (Seq. ID No. 8) |
| 11. 14 | 8 | AAT | CAT | TTT | CTG | ATG | CCG | AAG | (Seq. ID No. 9) |
| 2 | 9 | AAT | CAT | TTT | CTT | AAG | GGG | ACG | (Seq. ID No. 10) |
| 20 1 | 10 | ATT | CAT | TTT | TTG | CCG | CAG | AAT | (Seq. ID No. 11) |
| | 11 | ATT | CAT | $\mathbf{T}\mathbf{T}\mathbf{T}$ | CTT | CTT | TGG | CGT | (Seq. ID No. 12) |
| - F -A | 12 | AAT | CAT | TTT | CTG | ATT | AGG | AAG | (Seq. ID NO. 13) |
| | 13 | AAT | CAT | TTT | CTG | CCG | ACT | GCT | (Seq. ID No. 14) |
| | 1 | Asn | His | Phe | Leu | Ile | Lys | Pro | (Seq. ID No. 19) |
| 25 | 2 | Asn | His | Phe | Leu | Arg | Ser | Pro | (Seq. ID No. 20) |
| | 3 | Asn | His | Phe | Leu | Pro | Arg | \mathtt{Trp} | (Seq. ID No. 21) |
| | 4 | Asn | His | Phe | Leu | Pro | Lys | Val | (Seq. ID No. 22) |
| | 5 | Asn | His | Phe | Leu | Leu | Pro | Pro | (Seq. ID No. 23) |
| | 6 | Asn | His | Phe | Leu | Pro | Pro | Arg | (Seq. ID No. 24) |
| 30 . | 7 | Asn | His | Phe | Leu | Pro | Thr | Gly | (Seq. ID No. 25) |
| | 8 | Asn | His | Phe | Leu | Met | Pro | Lys | (Seq. ID No. 26) |
| | 9 | Asn | His | Phe | Leu | Lys | Gly | Thr | (Seq. ID No. 27) |
| | 10 | Asn | His | Phe | Leu | Pro | Gln | Asn | (Seq. ID No. 28) |
| | 11 | Asn | His | Phe | Leu | Leu | Trp | Arg | (Seq. ID No. 29) |
| | 12 | Asn | His | Phe | L u | Ile | Arg | Lys | (Seq. ID No. 30) |
| | 13 | Asn | His | Phe | Leu | Pro | Thr | Ala | (Seq. ID No. 31) |
| | | | | | | | | | |

For purposes of this application, discussions relating to the particular peptides will refer to the isolate numbers at the left side of the table. The Seq. ID No.'s relate to the computer-readable print-out which must be provided to the various patent offices.

To confirm that the <u>B. subtilis</u> peptides are tight-binding ligands, the following experiment was performed. 10⁷ phage of isolate #4 (NHFLPKV) (Seq. ID No. 15) and 10¹⁰ phage containing random 7-mer sequences were mixed with 10⁹ spores. This mixture was subjected to a single round of biopanning. The eluted phage were plaque-purified and genomic DNA was sequenced as described above. Seven of the ten phage examined contained the sequence of isolate #4. Thus, there was a 700-fold enrichment of this phage, clearly indicating that the isolate #4 peptide bound tightly to the spore.

Attempts to bind the spores of <u>B. subtilis</u> with the 4-mer peptide NHFL (Seq. ID No. 1) showed that sequence to be a poor ligand. However, the 5-mer sequence NHFLP (Seq. ID No. 39) showed tight binding.

In a search of the Swiss-Prot data base of characterized peptides for proteins containing the sequence NHFLP, seven proteins with this sequence were identified. Five eukaryotic proteins and two are B. subtilis proteins. first B. subtilis protein is SpsC (Database accession number BG10611), which contains the NHFLP sequence, near its amino terminus (i.e., MVQKRNHFLPYSLP-) (Seq. ID No. 16). SpsC appears to be involved in the synthesis of polysaccharides on the surface of the spore. It is probable that this protein uses its amino terminus to attach to a receptor on the spore surface. The instantly claimed peptide ligands may bind to the The second B. subtilis protein is UvrC (Database same site. accession number BG10349), an exonuclease involved in DNA The NHFLP sequence is found in the middle of UvrC, which contains 598 amino acids. Because UvrC is known to be cytoplasmic, a connection between this protein and the peptide ligands is not obvious.

Alternatively, differential display can be utilized to

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quickly find small molecule analogs or antagonists of present peptides (Greenwood, et al., Multiple display of foreign peptides а filamentous bacteriophage: Peptides plasmodium falciparium circumsporozoite protein as antigens. J. Mol. Biol. 206:821-827, 1991).

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Asn

Ser

His

Phe

(Seq. ID No. 26)

Leu Pro

There were only thirteen unique DNA sequences (out of a total of thirty) from B. subtilis spore-binding phage found. The frequency with which a particular sequence is found may directly reflect the tightness of binding of the encoded peptide. Although the sequences at positions five through seven are not identical, there is a clear preference for certain amino acids. Nearly one-third of all residues in positions 5, 6 and 7 are prolines (12/39), 31% are positively charged (5/39 Arg and 4/39 Lys), and the rest are hydrophobic or hydroxyl-containing. At position five, there is a strong preference for proline (6/13). Thus, it appears that these peptides bind to the same receptor on the spore coat.

The biopanning experiment described above was repeated using a library containing larger 12-mer peptides seen below:

Leu

Asn

Pro

Ala

Pro

Pro

^{1.} Asn His Phe Leu Lys Ser Gln Pro Gly Val Thr (Seq. ID No. 80) 2. Asn His Phe Leu Asn Pro Arg Ala Gln Ser Gln Val (Seq. ID No. 81) 3. His Asn Phe Leu Pro Pro Lys Met Gly Pro Thr (Seq. ID No. 82) Asp Phe 4. Asn Leu Pro Glu Pro Arg Leu Met Pro (Seq. ID No. \$3) 5. Asn His Phe Leu Ala Pro Gln Pro Pro Val Lys Pro (Seq. ID No. 84) Leu₃Met 6. Asn Pro Asn Pro Leu Leu Ala (Seq. ID No. &4) Met 7. Asn His Phe Leu_{2/}Ile Pro Pro Glu Pro Leu Arq Glu (Seq. ID No. 85) 8.

When the 12-mer peptides were compared with the 7-mer peptides, it appeared that no improvement occurred as a result of using the longer peptides.

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Selected peptides were analyzed for tight binding to the spore. Two peptides were synthesized initially: NHFLPKVGGGC (Seq. ID 16) and LFNKHVPGGGC (Seq. ID 17). The first has the amino-terminal sequence of peptide #4 plus a Gly₃ linker and a carboxy-terminal Cys. The second has a randomized sequence using the amino acids of peptide #4 plus the Gly₃ linker and carboxy-terminal Cys. The goal was to label these peptides at the carboxy-terminus with phycoerythrin and examine binding of test and control peptides by fluorescence microscopy and FACS sorting.

Initially, peptide-phycoerythrin conjugates were used for FACS. The advantage is that the conjugates are multivalent and the fluorescence characteristics are well suited for FACS. In some instances, peptides are first being reduced with tris(2-carboxyethyl)phosphine (TCEP) before conjugating with the phycoerythrin. Labeling with smaller fluorochromes such as monovalent 5-iodoacetamido-fluorescein is being used as an alternative.

In order to identify the receptor that interacts with the peptide, biotin-containing cross-linking agent that has been attached to a tight-binding peptide. Cross linkers examined sulfosuccinimidy1-2-[6-(biotinamido)-2-(pazidobenzamido)-hexanoamido]ethyl-1'c3'-dithiopropionate)sulfo-SBED). The molecule contains three different functional groups One arm consists of a biotin handle that can be used for purification using immobilized avidin. Another arm a sulfo-NHS (N-hydroxy-succinimido) provides amine coupling capability. When mixed with a tightbinding peptide, NHFLPKV plus GGGC (Seq. ID No. 99) extension, the cross linker is covalently coupled to the peptide through the ϵ -amino group of the carboxy-terminal lysine residue, with the release of N-hydroxy-succinimide. To assure coupling only through the ϵ -amino group of the lysine, the amino terminus of the peptide (i.e., the lpha-amino group of asparagine)

temporarily protected. The third arm contains a photosensitive phenyl azide that can be activated by exposure to UV light at wave lengths greater than 300 nm. The activated phenyl azide reacts with nucleophiles, especially amines, in the target molecule.

Once the peptide-cross-linker conjugate was prepared, it was mixed with spores for 10 minutes in the dark to allow peptide-receptor interaction. The complexes were exposed to UV (365 nm) light for 15 minutes at 0°C to allow cross-linking to the receptor. The spores were then collected by centrifugation, resuspended in SDS-PAGE loading dye (4% SDS, mercaptoethanol, 1 mM dithiothreitol, 125 nM Tris-HCl (pH 6.8), 10% glycerol and 0.05% bromophenol blue) and boiled for 8 minutes to solubilize spore coat proteins (including receptor) and to reduce the disulfide bond that attaches the peptide to the cross-linking agent. Intact spores were removed by centrifugation. The supernatant containing solubilized proteins was dialyzed (MW cutoff: 2000 Da) against phosphatebuffered saline (PBS). The sample was passed over a monomer avidin column and washed with PBS to remove proteins lacking a biotin-containing cross-link. The bound protein/receptor was eluted with PBS containing 2 mM biotin. The fraction containing eluted protein (measured by OD_{280}) was dialyzed against $\mathrm{H}_2\mathrm{O}$ and analyzed by SDS-PAGE. If one or more proteins were they were detected, analyzed by sequencing their terminus.

Biopanning with the heptamer phage display library was used to identify tight-binding peptides on the surface of <u>B</u> antracis spores. The spores were prepared from the avirulent delta-Ames strain of the organism (lacking the toxin-encoding plasmid pOX1) and were sterilized by gamma-irradiation by Diagnostics Systems Division of the U.S. Army Medical Research Institute of Infectious Disease, Fort Detrick, Maryland.

Four rounds of biopanning were performed in the manner described above. The genomic DNA of amplified eluates from each round were sequenced directly and genomic DNA of 27 single plaques from the fourth round amplified elute were also

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The DNA sequences from single plaques of the fourth round of biopanning are summarized in Table 2. Thirteen of the 27 sequences were the same as the dominant sequence found in the amplified eluate. Two other closely related sequences TYPIPIR (Seq. ID No. 41) and TYPIPFR (Seq. ID No. 42) were represented Another sequence TYPVPHR (SEQ. ID No. 43) three times each. similar to the previous two sequences was found once. The three last sequences define a tight binding sequence of the consensus formula $TYPX_1PX_2R$ wherein X defines hydrophobic residues and where the preferred X₁ is valine (V) or isoleucine (I) and X_2 is isoleucine (I), Phenylalanine (F) or Histidine A listing of the sequences is shown in Table 2. has been shown that the first four amino acids of the most common sequence (TSQN) (Seq. ID 44) is present in domain 3 of the B. anthracis protective antigen.

Sequences 4, 5, 6, and 9 are preferred sequences for binding the spore coat.

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TABLE 2: Nucleotide and Amino Acid Sequences from <u>B. anthracis</u> Spore Binding Phag

| | Isolate | | | | | | | | | | |
|-----------------------|---------|------------|------------|------------|------------|------------|------------|------------|------------------|------------------|------|
| 5 | 1 (1) | AAT Asn | AGT Ser | GTT Val | ACT Thr | CTT Leu | GAG Glu | CCG Pro | (Seq I (Seq I | D No. | |
| | 2 (1) | AAG Lys | CCG Pro | AGG Arg | CAG Gln | CCG Pro | GGT Gly | TTG Leu | (Seq. : | ID No. ID No. | |
| 10 | 3(1) | TCT Ser | ACT Thr | CCG Pro | GCG Ala | TGG Trp | CTG Leu | TCG Ser | (Seq. | ID No. ID No. | |
| | 4 (13) | ACT Thr | AGT Ser | CAG Gln | AAT Asn | GTG Val | CGG Arg | ACG Thr | (Seq. | ID No. ID No. | |
| | 5 (3) | ACT Thr | TAT Tyr | CCT Pro | ATT Ile | CCG Pro | ATT Ile | CGT Arg | (Seq. | ID No. ID No. | |
| © 15 √0 | 6 (3) | ACT Thr | TAT Tyr | CCT Pro | ATT Ile | CCG Pro | TTT Phe | CGT Arg | (Seq. : | ID No. ID No. | |
| 20 mg fill mg fill mg | 7 (1) | TCT Ser | TAT Tyr | CCT Pro | CAT His | GGT Gly | CAG Gln | ATT Ile | (Seq. : | | |
|) [20 | 8 (1) | TTT Phe | ACT Thr | GGG Gly | ACT Thr | CTT Leu | AAT Asn | CCT Pro | (Seq. : | ID No. ID No. | |
| | 9 (1) | ACT Thr | TAT Tyr | CCG Pro | GTG Val | CCG Pro | CAT His | CGG Arg | (Seq. | | |
| } ⊸ | 10 (1) | CGG Arg | ACT Thr | CCT Pro | TCG Ser | CTT Leu | CCT Ser | AGT Pro | (Seq.] | | |
| 1 25 1 25 | 11 (1) | TTT Phe | AGT Ser | GTT Val | CCT Pro | CGT Arg | ATG Met | CCG Pro | (Seq.] | | |
| ₩ | The | number | : in (|) re: | fers t | to the | numb | er of | phage c | ontair | nina |

The number in () refers to the number of phage containing the sequence.

Studies with <u>B cereus T</u> were undertaken using methods described above. After the fourth round of biopanning, individual phages were cloned. Twenty-two phage were picked, the genomic DNA prepared from each, and the DNA coding regions sequenced. The results revealed 8 unique DNA sequences. (See Table 3.) A heptapeptide VTSRGNV (Seq. ID No. 100) having tight-binding properties was identified. This sequence emerged from the pooled genome sequence of amplified phage following the third round of biopanning. The following unique DNA sequences were identified:

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|-----------|--------|-------|------------|------------|------------|------------|------------|------------|------------|--------|------------------|--|
| | But CI | TABLE | 3. | B. ce | ereus | T spo | or t | ight- | bindi | ng pep | tides: | |
| | J _1. | (2) | ACG Thr | CAT His | CGT Arg | TTG Leu | CCT Pro | TCT Ser | CGG Arg | | ID No. ID No. | |
| 5 | 2. | (13) | GTT Val | ACT Thr | AGT Ser | AGG Arg | GGG Gly | AAT Asn | GTT Val | | ID No. ID No. | |
| | 3. | | AAG Lys | CTG Leu | TGG Trp | GTG Val | ATT Ile | CCT Pro | CAG Gln | | ID No. ID No. | |
| 10 | 4. | | TAT Tyr | TCG Ser | CCT Pro | CCT Leu | CAT His | AGG Arg | CAT His | | ID No. ID No. | |
| | 5. | | TCG Ser | TAT Tyr | CCT Pro | CCG Pro | TAT Tyr | TTT Phe | GAT Asp | | ID No. | |
| | 6. | (2) | CTT Leu | TTG Leu | TCG Ser | CCT Pro | CTG Leu | CAT His | CGT Arg | | ID No. ID No. | |
| 15] | 7. | | TTT Phe | GAT Asp | TCT Ser | CCG Pro | CTT Leu | CGT Arg | CGG Arg | | ID No. ID No. | |
| | 8. | | TGG Trp | TCG Ser | CCG Pro | CTG Leu | CAT His | AAG Lys | CAT His | | ID No. ID No. | |
| 20 10. | | | | | | | | 13 | | | ~ | |

One DNA sequence (Seq. ID No. (111)), found in 13 of the 22 phage, encoded the previously identified tight-binding peptide VTSRGNV. A second sequence was obtained from an inspection of the remaining 7 unique phage DNA sequences. Four of these sequences (4, 6, 7 and 8) contained all or most of the closely related sequence SPL(H or R) (R or K)H. Such results are highly suggestive of a true tight-binding peptide sequence.

A competitive biopanning study was performed using phage displaying unique peptide sequence 8 (WSPLHKH) (Seq. ID No. 117) in this study. 10¹⁰ phage from a random phage library and 10⁷ phage displaying sequence #8 were mixed together, and one round of biopanning was performed using spores of B cereus T. The eluted phage were plaque-purified and genomic DNA was sequenced for twenty phage. Six of the twenty phage contained the sequences of isolate #8. Thus, there was a 300-fold enrichment of this phage, indicating tight binding.

In efforts to find additional tight-binding peptides the wash buffer or wash conditions have been systematically modified. One alteration is the use of either 0.5% or 0.01% Tween 20 and either 3 or 10 washes. These conditions were used in biopanning for <u>B subtilis</u>, wherein number of washes were

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reduced from 10 to 3. After four rounds of biopanning, an amplified eluted phage pool was sequenced. The results indicate that by changing certain parameters, it is possible to detect new tight binders for some spore species.

It was found that it was possible to precipitate the amplified phage for 30 minutes instead of overnight. Using the abbreviated method, it was possible to omit all titering of phage between the rounds. Using this method, an approximate concentration of amplified phage (1.75 X 10¹³ pfu/ml) is assumed. Omitting these steps allows a four-round biopanning experiment to be completed in two 12-hour days without affecting results.

Modified versions of the biopanning procedure can also be used wherein phage are permitted to bind spores. Binding complexes are recovered by centrifugation. Complexes are mixed with <u>E. coli</u> to permit phage amplification (under conditions where <u>B. subtilis</u> growth is inhibited), and amplified phage are subjected to additional rounds of biopanning. Tight-binding phage are then recovered by centrifuging spores plus bound phage through a density gradient.

The propensity of the peptides provided herein to bind to spore surfaces makes it possible to capture and identify target bacteria and spores to which the peptides bind. For example, when tagged sequences which bind to the surface of spores of B. subtilis, particularly 5-mer to 12-mer sequences, are placed in the environment believed to contain B. subtilis spores, the presence of the bacteria of interest are identified. Tags such as fluorescent, phosphorescent or colorimetric tags make it possible to visualize the presence of the bacteria. tags, such as radioactive tags, may require other equipment such as scintillators to determine the presence or absence of the target organisms. The method described above is particularly useful for identifying contamination of water and food that might cause disease when ingested. Contamination of the air might be established using methods of the invention. latter is particularly important when the possible contaminant is B. anthracis. It would be possible to attach the peptides

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identified as having the appropriate binding properties to solid supports to capture spores or spore-forming organisms which bind to the peptide. The particular support will depend on the use. For example, appropriate supports may be natural fibers or polymers which may be in the form of filtering devices, tapes or sponges. Supports having the binding peptides may be used as protective barriers such as masks.

Purified peptides formulated in pharmaceutically acceptable carriers such as buffered saline may be administered to animals in an appropriate amount to elicit an immune response or to bind to the spore to cause alteration in pathogenicity. The method of administration will depend on the organism and the site of infection. Formulations for inhalation may also be buffered to prevent damage to tissue.

Polyclonal antibodies to the sequences of interest can be produced in animals and purified directly from the spleen It is also common to isolate spleen cells from the cells. animal for purposes of producing antibodies. These cells can then be fused with an immortal cell line and screened for monoclonal antibody secretion. Purified antibodies that specifically bind the peptide are within the scope of the present invention. The antibody can be labeled by means generally known in the art using, for example, fluorescent, radioactive or phosphorescent markers, or tags may used in conjunction with a labeled secondary antibody in methods such as ELISA tests. Monovalent, divalent or single chain antibodies can be made which bind the peptides of the invention.

Anti-idiotype antibodies can also be made by means commonly known in the art. Antibodies to the present peptides can exhibit idiotypic mimicry and can be administered to provide protection against bacterial infection. Antibodies to the spore-binding peptides provided herein can be administered to susceptible hosts to block SpsC binding to the spore surface, thus inhibiting development of clinical disease.

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